Endothelin-1 Is a Critical Mediator of Myogenic Tone in Tumor Arterioles: Implications for Cancer Treatment

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ABSTRACT

Although derived from the host tissue, the tumor vasculature is under the influence of the tumor microenvironment and needs to adapt to the resistance to blood flow inherent to the dynamics of tumor growth. Such vascular remodeling can offer selective targets to pharmacologically modulate tumor perfusion and thereby improve the efficacy of conventional anticancer treatments. Radiotherapy and chemotherapy can, indeed, take advantage of a better tumor oxygenation and drug delivery, respectively, both partly dependent on the tumor blood supply.

Here, we showed that isolated tumor arterioles mounted in a pressure myograph have the ability, contrary to size-matched healthy arterioles, to contract in response to a transmural pressure increase. This myogenic tone was exquisitely dependent on the endothelin-1 pathway because it was completely abolished by the selective endothelin receptor A (ETα) antagonist BQ123. This selectivity was additionally supported by the large increase in endothelin-1 abundance in tumors and the higher density of the ETα receptors in tumor vessels. We also documented by using laser Doppler microprobes and imaging that administration of the ETα antagonist led to a significant increase in tumor blood flow, whereas the perfusion in control healthy tissue was not altered. Finally, we provided evidence that acute administration of the ETα antagonist could significantly stimulate tumor oxygenation, as determined by electron paramagnetic resonance oximetry, and increase the efficacy of low-dose, clinically relevant fractionated radiotherapy.

Thus, blocking the tumor-selective increase in the vascular endothelin-1/ETα pathway led us to unravel an important reserve of vasorelaxation that can be exploited to selectively increase tumor response to radiotherapy.

INTRODUCTION

The tumor vasculature has unique features, which may be viewed as many selective targets for anticancer strategies. Accordingly, the last 15 years have witnessed the development of drugs targeting tumor vessels, namely the angiogenic (1, 2) and antiangiogenic agents (3). Although the goals of these approaches are different (e.g., to prevent new blood vessel formation and to occlude/destroy pre-existing vessels, respectively), they both exploit the specific phenotype of endothelial cells lining tumor blood vessels to selectively target tumor perfusion (4). An interest also simultaneously developed for opposite strategies aiming to enhance the tumor perfusion (5, 6). The rationale for such qualitative improvement in the function of tumor blood vessels is to increase the efficacy of radio- or chemotherapy (7, 8). This concept implies that the effects on tumor blood flow need to be transient and selective. Although the action of vasomodulatory drugs can be controlled by adapting their protocol of administration, the selectivity issue is limiting and matter of investigation (9).

We and others have reported that drugs like nitric oxide donors (10, 11), nicotinamide (12), bradykinin agonist (13), and insulin (14), but also radiations and chemotherapeutic agents themselves (15–17), could modulate tumor blood flow, oxygenation, and permeability, thereby optimizing associated antitumor treatments. The validity of these adjuvant approaches to conventional chemo- and radiotherapy was usually verified a posteriori by documenting a better tumor response with very limited insights on the mechanisms of the tumor selectivity. Conversely, studies that did not look for a gain in treatment efficacy have identified the existence of tumor-specific vascular reactivity to various products including endothelin (18), endothelin agonist (19), angiotensin (20, 21), noradrenaline (22), hemoglobin A (23), and tumor necrosis factor α (24).

We reasoned that the identification of vasomodulatory pathways truly selective of the tumor vasculature could arise from the ex vivo comparison of the intrinsic pharmacological reactivity of vessels issued either from tumors or from the tumor-hosting tissues. Indeed, the variety of chemokines and cytokines released by tumor cells are known to trigger regulatory processes within the tumor vasculature leading to phenotypic shift including escape from immunosurveillance and adaptation to hypoxia (2, 25, 26). Therefore, it was very likely that the tumor microenvironment could also, directly or indirectly, induce profound alterations in the tumor vessel reactivity. Accordingly, using a myograph (originally dedicated to study the physiopathology of normal arterioles), we reported recently that the acetylcholine-induced nitric oxide-mediated vasodilation was dramatically altered in tumor arterioles (15). Also, in the same study, we documented how ionizing radiations by reversing the lack of NO-mediated response of tumor vessels could increase the sensitivity of the tumor to further X-ray fractions (15).

In the current study, our reflection started from the growing evidence that endothelin-1 production is increased in many tumor types where it plays crucial roles in proliferation and angiogenesis (27–29). In these studies, however, there was no mention of the powerful vasoconstrictive effects of endothelin-1 (ET-1) and its potential action on the contractile tone of tumor vessels. Indeed, although ET-1 can induce vasodilation through activation of the endothelin receptor B (ETβ), a powerful vasoconstriction is obtained when ET-1 interacts with the endothelin receptors A and B (ETα and ETβ) located on smooth muscle cells. In this study, we therefore asked ourselves whether the tumor production of ET-1 could influence local vascular tone and whether blocking ET-1 signaling could have an impact on tumor blood flow and thereby be exploited to promote conventional antitumor therapy.

MATERIALS AND METHODS

Mice, NMRI, C57BL/6, and C3H/He male mice (Elevage Janvier, Le Genest-St-Ise, France) were used in experiments with transplantable liver tumor (TLT; mouse hepatocarcinoma; Ref. 30), Lewis Lung carcinoma (mouse lung carcinoma; Ref. 31), and Fibrosarcoma-II (mouse fibrosarcoma; Ref. 32) syngeneic tumor cells, respectively. Anesthetized mice (ketamine/xylazine) received i.m. injections of 103–105 tumor cells in the vicinity of the saphenous arteriole of the posterior right leg. The pre-existing arteriole was progressively co-opted as the

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Note: P. Sonveaux and C. Dessy contributed equally to this work.

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tumor grew (see Fig. 1G). The tumor diameters were tracked with an electronic caliper. When the tumor diameter reached 4.0 ± 0.5 mm, mice were sacrificed or randomly assigned to a treatment group. When indicated, they received an i.p. injection of the selective ET_{A} antagonist BQ123 (0.5, 1, and 2 mg/kg; Sigma, Bornem, Belgium) or saline alone. Each procedure was approved by the local authorities according to national animal care regulations.

**Immunoblottings and Immunostainings.** Immunoblottings were carried out, as described previously (15, 33), with a sheep polyclonal antibody against ET_{A} (Abcam, Cambridge, United Kingdom) or ET_{B} (Alomone Labs, Jerusalem, Israel). For immunostainings, collected tissue samples were cryosliced and probed with a rabbit polyclonal antibody against ET-1 (gift from Ronald van Beneden, University of Louvain, Brussels, Belgium; Ref. 34). Endogenous peroxidase activity was inhibited by 0.3% H_2O_2 in PBS, and Envision system (Dako, Glostrup, Denmark) was used for revelation. Sections were finally counterstained with Mayer’s hematoxylin. Quantitative image analyses of immunostained arteriole slices (5 arterioles/condition and 3 sections/animal) were performed with AnalySIS software (Soft Imaging System, Münster, Germany).

**Myograph Assays.** Tumor saphenous arterioles and size-matched arterioles from healthy mice were dissected under a stereoscopic microscope and mounted on a 110P pressure myograph (DMT, Aarhus, Denmark). No macroscopic differences were detectable between the tumor and control arterioles (diameters of 284 ± 14 and 299 ± 17 μm, respectively; P > 0.1). Changes in the outer diameters were tracked and measured with the Myoview software (DMT). To evaluate the myogenic tone, isolated arterioles were left to recover at physiological pressure for 45–60 min in no-flow condition (20 mm Hg; 37.5°C). Then, the pressure was increased by steps of 20 mm Hg (that were maintained for 15 min to allow arteriole response). For each pressure step, the active vessel diameter was determined after a 60-min preincubation in the calcium-containing physiological salt solution buffer with BQ123 (1 μM).

For the establishment of the ET-1 dose-response curve, isolated arterioles were left to recover at physiological pressure for 45–60 min in no-flow condition in physiological salt solution medium (60 mm Hg, 37.5°C); additive doses of ET-1 (Sigma) were then delivered in the bathing medium. For each vessel used in this study, the ability of the vessels to contract upon application of a depolarizing KCl solution was verified at the end of the experiment and compared with a similar contraction performed at the very beginning of the experiment. If these two contractions differed by >10%, the experiment was disregarded.

**Tumor Blood Flow Monitoring.** Tumor perfusion was measured with a Laser Doppler imager (Moor Instruments) and with Laser Doppler microprobes (Oxyflo; Oxford Optronix). Briefly, for the Laser Doppler imaging measurements, mice were anesthetized and fur was removed from the limbs using a depilatory cream. They were placed on a heating pad (37°C) to minimize variations in temperature. The average perfusions of the tumor-bearing leg and of the control leg were evaluated on the basis of colored histogram pixels. For the Oxyflo measurements, fiber-optic microprobes (Laser Doppler + thermocouple) were introduced into the tumor. Back scattering measurements were used to validate the absence of movement artifacts. A 10-min baseline of stable recordings was obtained before treatment administration through the catheterized tail vein; data were collected continuously at a sampling frequency of 20 Hz.

**Electron Paramagnetic Resonance (EPR) Oximetry.** This real-time, O_2- nonconsuming technique was used to track the changes in pO_2 induced by BQ123 administration. EPR oximetry relies on the oxygen-dependent broadening of the EPR linewidth of a paramagnetic oxygen sensor preimplanted in the tumor. Accordingly, 50 μl of a suspension (100 mg/ml) of the O_2-sensitive probe (charcoal wood powder, CX0670–1; EM Science, Gibbstown, NJ) were injected in the center of the tumor 24 h before X-ray irradiation, as described previously and validated (10, 14, 15). The tumors of anesthetized mice were placed in the center of the surface coil. EPR spectra were recorded using an

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**Fig. 1.** Immunodetection of endothelin-1 (ET-1) in tumor and tumor vessels. ET-1 antibodies were used to immunostain (A) cardiac tissue from healthy mice (cl indicates the lumen of a coronary artery); (B) transplantable liver tumor; (C) Lewis Lung carcinoma tumor; and (D) Fibrosarcoma-II tumor. —— delineate the healthy muscular tissue (m) surrounding tumors or the central necrotic core (nc). Bar = 100 μm. Saphenous arterioles from (E) healthy or (F) tumor-bearing mice were also immunostained for ET-1 expression. G, shown is the anatomical location of the saphenous arteriole located between the saphenous vein (v) and nerve (n); the —— shows the limits of the tumor and the arrowhead indicates where the arteriole enters the tumor (e.g., where arterial segments were isolated for myograph studies).
Results

Expression of ET-1 in Tumor Cells and in the Tumor Vasculature. We first characterized the ET-1 production in 3 syngeneic tumor models (TLT, Lewis Lung carcinoma, and Fibrosarcoma-II) obtained by i.m. injections of tumor cells. When reaching a diameter of 4.0 ± 0.5 mm, the tumors were resected and processed for immunostaining with ET-1 antibodies; mouse heart was used as control for antibody specificity (see Fig. 1A). As shown in Fig. 1, B–D, the different tumors were found to produce large amounts of ET-1 that were detectable both inside the tumor cells and in the extracellular space. Very low amounts of ET-1 were found in the healthy surrounding muscular tissues (see Fig. 1, E and C) and in the necrotic core of the tumor (see Fig. 1D). Importantly, the co-opted tumor vasculature was strongly labeled, revealing a 13-fold increase (P < 0.05; n = 15) in ET-1 in the vessel wall when compared with control size-matched arterioles (Fig. 1, E and F).

Differential Reactivity of Tumor Arterioles to ET-1. To evaluate the vasomodulatory effects of the increased ET-1 concentrations in tumor arterioles, we isolated TLT-co-opted arterioles (see Fig. 1G) and size-matched arterioles from healthy animals. These arterioles were mounted on a pressure myograph and, after pressurization and equilibration, exposed to increasing doses of ET-1. Results are expressed as percentage of the maximal contraction obtained with 10⁻⁶ M ET-1. Sigmoidal fitting shows that the dose/Effect curves for healthy and tumor vessels (R² = 0.91 and 0.98, respectively) were significantly different (P < 0.01; n = 5); note that some SE are smaller than symbols; bars, ± SE; B, pools of arterioles from healthy and transplantable liver tumor-bearing mice were lyzed (to obtain enough material) and were immunoblotted with endothelin receptor A (ETA) and endothelin receptor B (ETB) antibodies. Shown are representative immunoblots and a bar graph depicting the densitometric analysis of two different immunoblots (2 pools of 10 arterioles/condition).

ETA inhibition triggers a tumor-specific increase in blood flow and oxygenation. We then investigated the implication of the ET-1/ETA pathway in the myogenic tone of TLT tumor-co-opted arterioles by using the specific ETA antagonist BQ123. The effect of the preincubation with BQ123 was dramatic, because it completely abrogated the myogenic tone (Fig. 3B; P < 0.01). In fact, the active diameter curve under ETA inhibition (Fig. 3B, black triangles) was not different from the passive diameter curve (Fig. 3B, open circles). As a control, we also demonstrated that ETA inhibition left unaffected the active diameter of size-matched arterioles from healthy mice (Fig. 3A).

ETA Inhibition Triggers a Tumor-Specific Increase in Blood Flow and Oxygenation. We then investigated whether the BQ123-mediated abrogation of the myogenic tone (as elicited ex vivo) was correlated with an increased tumor perfusion in vivo. Accordingly, TLT tumor-bearing mice received a bolus i.p. injection of 1 mg/kg BQ123 or saline, and the perfusion of both the tumor-bearing and the control legs was monitored using a Laser Doppler imaging system. In BQ123-treated mice, we consistently observed an important increase in the blood flow of the tumor-bearing leg (Fig. 4A, top panel, white circle), whereas the healthy opposite leg failed to show any significant changes in perfusion (Fig. 4A, top panel, white arrows). Importantly, isovolemic saline i.p. injection barely increased the basal perfusion in the tumor-bearing leg (see Fig. 4A, lower panels, and Fig. 4B). We also used Laser Doppler needle probes directly introduced in the tumors to validate the results.
Fig. 3. Tumor arterioles exhibit an endothelin receptor A-dependent myogenic tone. Isolated arterioles were mounted on a pressure myograph and allowed to respond to a step-by-step increase of the luminal pressure. Changes in vessel diameter were tracked in bathing media containing calcium (active diameter, ▲; n = 10) or not (passive diameter, ○; n = 10). Saphenous arterioles were dissected from healthy mice (A) or were isolated from transplantable liver tumor (after co-option; B). In some experiments, arterioles were preincubated with the endothelin receptor A antagonist BQ123 (1 µM) in the calcium-containing medium (active diameter, ▲; n = 3); ns, not significantly different; **, P < 0.01 between the indicated dose/effect curves; bars, ±SD.

obtained with the imager. A similar increase in blood flow was observed after BQ123 administration (Fig. 4C).

Using EPR oximetry, we also evaluated the BQ123-induced changes in tumor pO2. Fig. 4D shows that 60 min after administration of the ETA antagonist (but not of the saline solution), the tumor oxygenation was significantly increased over basal level (+174%; P < 0.01; n = 7). Of note, the measured pO2 corresponds to the mean value in a tumor volume of ± 10 mm3 (i.e., the volume of charcoal dispersion as determined by histology) and, therefore, does not preclude more dramatic variations in specific regions of the tumor.

ETA Inhibition Radiosensitizes Tumors. In various tumor models, improved tumor perfusion is associated with a net increase in tumor oxygenation, which may be exploited to increase the effectiveness of radiotherapy (10, 14, 15). Therefore, we tested the hypothesis that ETA inhibition could radiosensitize tumors. Accordingly, we daily determined the tumor diameters of mice receiving (or not) fractionated radiotherapy, and evaluated the effects of BQ123 administered 45 min before each radiation fraction (to have the maximum increase in pO2 at the time of X-ray exposure; see Fig. 4). As shown in Fig. 5, A and B, and Table 1, when fractionated radiotherapy (5 × 2 Gy and 2 × 6 Gy) was combined to the BQ123 pretreatment, the overall benefit on tumor growth retardation was higher than the sum of the effects of the two treatments administered separately (P < 0.01; n = 8–11). Similar data were obtained with two other tumor models, namely Lewis Lung carcinoma tumors and Fibrosarcoma-II (data not shown); both tumor types overexpress ET-1 (see Fig. 1, C and D) and arterioles issued from the former were shown to also exhibit myogenic tone (data not shown).

A dose-response curve was also generated using three different doses of BQ123 (0.5, 1, and 2 mg/kg) in combination with the 2 × 6 Gy protocol (Fig. 5C). Although limitations in BQ123 availability precluded additional exploration, it is noteworthy that with the higher dose (2 mg/kg), two mice (of 6) were in remission 5 weeks after treatment (see Fig. 5 legend). Of note, at each dose used in this study, BQ123 (two or five i.p. injections) only slightly restrained tumor growth by itself (see Fig. 5, A and B, and Table 1).

DISCUSSION

The two major findings of this study are: (a) that basal production of ET-1 in tumors leads to the development of a myogenic tone that represents an important reserve for vasorelaxation; and (b) that the use
were determined (condition, three doses of BQ123 were tested (0.5, 1, and 2 mg/kg), and mean tumor doubling times were determined (P < 0.03; n = 6–8). Of note, the beneficial effect of the 2 mg/kg dose is underestimated because the doubling time was not determined for 2 mice (of 6), which were in complete remission 5 weeks after the initiation of treatment. BQ123 alone (at each dose tested) failed to produce remission 5 weeks after the initiation of treatment. When coadministered with radiotherapy, BQ123 (or vehicle) was delivered i.p. 45 min before each irradiation; bars ±SD.

Table 1 Mean regrowth delays for TLTa tumors as a function of ionizing radiation exposure(s) and BQ123 treatment(s)

<table>
<thead>
<tr>
<th>BQ123 (mg/kg)</th>
<th>X-rays (# fraction × dose)</th>
<th>Tumor doubling time (days)</th>
<th>Mean regrowth delay (days)</th>
<th>Mice (n)</th>
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</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>4.1 ± 0.7</td>
<td>—</td>
<td>11</td>
</tr>
<tr>
<td>0</td>
<td>5 × 2 Gy</td>
<td>8.5 ± 0.6</td>
<td>4.4 ± 0.8</td>
<td>11</td>
</tr>
<tr>
<td>+</td>
<td>5 × 0 Gy</td>
<td>4.6 ± 0.3</td>
<td>0.5 ± 0.7</td>
<td>11</td>
</tr>
<tr>
<td>+</td>
<td>5 × 2 Gy</td>
<td>13.6 ± 0.7</td>
<td>9.5 ± 0.8</td>
<td>11</td>
</tr>
<tr>
<td>+</td>
<td>2 × 6 Gy</td>
<td>13.2 ± 0.9</td>
<td>9.1 ± 0.9*</td>
<td>11</td>
</tr>
<tr>
<td>+</td>
<td>2 × 0 Gy</td>
<td>4.6 ± 0.4</td>
<td>0.5 ± 0.8</td>
<td>11</td>
</tr>
<tr>
<td>+</td>
<td>2 × 6 Gy</td>
<td>23.0 ± 1.4</td>
<td>18.9 ± 0.8*</td>
<td>8</td>
</tr>
</tbody>
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TLT, transplantable liver tumor.

a P < 0.01, versus additive effects of BQ123 and X-ray administered separately.

of an ETA antagonist can selectively promote tumor perfusion and oxygenation, and consequently increase the effectiveness of tumor radiotherapy.

The existence of a myogenic tone is almost impossible to quantitatively appreciate in vivo, because it requires the tracking of changes in vascular resistance in response to an increase in transmural pressure. Here, we used a unique experimental model specifically dedicated to this purpose, e.g., microdissected tumor vessels mounted in a myograph allowing step increases in pressure. This ex vivo mode of detection should not mask the in vivo importance of the myogenic tone in the autoregulation of blood flow by arterioles. Indeed, in normal vascular beds, spontaneous (oscillating) and pressure-induced variations in the muscular tone, known as vasomotion (36) and myogenic tone (35, 37), respectively, offer to the tissue circulation the intrinsic ability to maintain a relatively constant blood flow despite variations in blood pressure. In the case of tumors, such autoregulatory mechanisms are very likely to be needed to face the constant remodeling of tumor vascularization. Here, we have identified the implication of ET-1 as a trigger of the myogenic tone of tumor vessels. The signal transduction appears to transit through the smooth muscle-specific ETA receptor (the abundance of which is increased by a muscle-specific ETA receptor (the abundance of which is increased by...completely abrogated the pressure-dependent development of vascular tone by tumor arterioles.

That the implication of the ET-1/ETA pathway in the development of a myogenic tone is specific of the tumor vascular bed was additionally authenticated by the increase in blood flow observed after treatment with BQ123 in tumors but not in control tissue from the same mouse (see Fig. 4). Interestingly, ET-1 has been shown to exert its vasconstrictive effects when applied from the adventitial side of the vessels (38, 39), which in the case of ET-1-producing tumors is very likely to account for local (specific) effects. Moreover, we found an increased vascular ETA expression in tumor arterioles, which is also very likely to contribute to the observed myogenic tone. Exaggerated production of ET-1 has been reported previously to mediate similar increase in myogenic constriction in cerebral arteries exposed to oxidized low-density lipoprotein (40) and arterioles isolated from hypertensive rats (41). Together with our study, these reports indicate that pathological states such as atherosclerosis, hypertension, and tumor growth can transform vessels devoid of myogenic tone in vessels prone to blood flow autoregulation. Of note, the phenotype shift can be considered as yet more dramatic for tumor vessels, because we used vessels with an external diameter of 300 μm, which is higher than the usually accepted diameter for resistance arterioles (<170 μm).

The observed ex vivo and in vivo vasodilating effects of BQ123 imply that functional smooth muscle cells are the target of this drug in tumor vessels. The presence of smooth muscle cells or mural cells on vessels is viewed as a hallmark of maturation (42), and it is generally accepted that month/year-old human tumors are more likely to recruit pericytes on a larger proportion of their vasculature (versus rapidly growing mouse tumors). This suggests that functional tumor vessels constitute a yet more attractive target for selective vasomodulatory treatments in humans. Although additional studies are required to evaluate how our data can be extrapolated in the clinics, the tumor vessel selectivity of ETA antagonist treatment, as identified in mouse, offers strong bases for the establishment of an adjuvant strategy aiming to increase the effectiveness of conventional antitumor treat-
mments (by acting on tumor blood flow and oxygenation). Adequately scheduled combination of these latter approaches with antiangiogenic drugs certainly represents a very attractive area of investigation for future cancer therapy development.

In this study, we provide evidence that acute administration of BQ123 (i.e., i.p. injection 45 min before local tumor irradiation) exerts profound effects on the efficacy of low-dose radiotherapy (5 × 2 Gy and 2 × 6 Gy). Importantly, these effects were shown to be ETA antagonist dose-dependent and obtained using clinically relevant schemes of fractionated radiation administration; a complete remis-

sion was even obtained in mice treated with the highest BQ123 dose tested in this study. Although a small effect of BQ123 alone was observed in the absence of irradiation (probably attributable to the antiproliferative effect of this drug; Refs. 27–29), the combination of both irradiation and the ETA antagonist led to a synergistic effect in both irradiation protocols, increasing the tumor regrowth delay by >50% over the additive effects (see Table 1). We additionally documented that the radiosensitizing effects of BQ123 were very likely to arise from the increase in tumor oxygenation directly associated with the elevation in tumor blood flow (see Fig. 4).

The growing amounts of reports documenting the increase in ET-1 expression in various human tumors (reviewed in Ref. 29) indicate that this approach of blocking the ETA receptor opens very interesting perspectives in the treatment of many tumor types. Moreover, although we have here only documented the impact of modulating the ET-1/ET(A) transduction pathway for radiotherapy, it can be postulated that chemotherapy could also take advantage of a better tumor perfusion and drug delivery. Importantly, this study also confirmed the potential of the myograph assay to identify the existence of a differential reactivity in arterioles issued from animal tumors characterized by different environments (see also Refs. 15, 33). This ex vivo pharmacological evaluation of tumor vessels could easily be extended to vessels isolated from human tumor biopsies, and thereby could lead to new insights on “provascular” adjuvant treatments susceptible to improve conventional anticancer treatments.

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